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## Structural determinants of Sleeping Beauty transposase activity.

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## ABSTRACT

Transposases are important tools in genome engineering, and there is considerable interest in engineering more efficient **ones**. Here we seek to understand the factors determining their activity using the Sleeping Beauty transposase. Recent work suggests that protein co-evolutionary information can be used to classify groups of physically connected, co-evolving residues into elements called 'sectors', which have proven useful for understanding the folding, allosteric interactions, and enzymatic activity of proteins. Using extensive mutagenesis data, protein modeling and analysis of folding energies, we **show** that 1) The Sleeping Beauty transposase contains two sectors, which span across conserved domains, and are enriched in DNA-binding residues, indicating that the DNA binding and endonuclease functions of the transposase coevolve; 2) Sector residues are highly sensitive to mutations, **and most** mutations of these residues strongly reduce transposition rate; 3) Mutations with a strong effect on free energy of folding in the DDE domain of the transposase significantly reduce transposition rate. 4) Mutations that influence DNA and protein-protein interactions generally reduce transposition rate, although most hyperactive mutants are also located on the protein surface, including residues with protein-protein interactions. This suggests that hyperactivity results from the modification of protein interactions, rather than the stabilization of protein fold.

## INTRODUCTION

Recent findings identified a structural organization of protein domains that is distinct from their known hierarchical organization into secondary and tertiary structural elements. These structures, termed 'sectors'<sup>1</sup> form physically connected networks of coevolving residues within proteins, and span across secondary structural elements. Sectors are identified using multiple alignments with a procedure called Statistical Coupling Analysis (SCA), which uses the covariance matrix of amino acid variability at different positions of the alignment, and their conservation<sup>1</sup>. It has been noticed that the residues that show correlated evolution in the alignments have a block structure in the SCA matrix: they can be partitioned into clusters of residues, which show correlated evolution within the cluster, but are essentially uncorrelated with residues of other clusters. These groups of coevolving residues were termed 'sectors', in analogy to financial sectors<sup>1,2</sup>. Several important biological properties of proteins are determined by sectors: although they typically make up only 10-30% of the residues of a protein, they were shown to significantly contribute to the specification of protein folds<sup>3</sup>, allosteric communication in proteins<sup>4</sup>, and evolution of novel functions<sup>5</sup>. Since it is possible to engineer functional artificial protein folds based purely on sector information<sup>6</sup>, or modify their functions using sector residues<sup>5</sup> (at least in small domains), sectors are of considerable importance also for protein engineering. However, most work to date on the architecture of sectors, functions and importance of sectors have focused on relatively few single-domain proteins, often with only a single sector<sup>1,4,5,7</sup>, and the number of studies with multidomain and multisector proteins is low<sup>1,8</sup>. Thus, it is unclear to what degree the current findings can be generalized, and whether sectors are of similar importance in more complex multi-domain structures as in small proteins<sup>2</sup>.

Most DNA transposons contain a single gene encoding the transposase protein, which is flanked by terminal inverted repeats (TIRs). Transposons 'jump' by a cut-and-paste mechanism, during which the transposase moves the sequence flanked by TIRs to a new genomic location. Since transposases require only the TIRs, and any sequence flanked by TIRs can be moved by externally supplied transposases, they can be used for gene transfer<sup>9</sup>. In consequence, transposons are popular tools that are widely used for genome engineering, including cancer gene identification by insertional mutagenesis<sup>10</sup>, germline transgenesis<sup>11</sup>, somatic gene transfer for gene therapy<sup>9</sup>, or cellular reprogramming<sup>12</sup>. Their primary advantage over viral vectors for gene therapy is that they have considerably fewer side effects, including

low immunogenicity and genotoxicity, while, at least for some applications, they provide stable transgene expression levels with efficiency matching viral vectors<sup>9</sup>. Several transposon systems are currently applied as genome engineering tools, including the piggyBac, Tol2 and Sleeping Beauty transposons<sup>13–18</sup>. The first DNA transposon tool capable for gene transfer in vertebrates was Sleeping Beauty (SB), which was reconstructed from extinct Tc1/mariner transposons in fish<sup>19</sup>. Sleeping Beauty, and especially its hyperactive variant<sup>20</sup> is still one of the most widely used transposon tool, and it is the only transposon vector being currently in human clinical trials<sup>21,22</sup>.

In this work, using our extensive mutagenesis data available for the Sleeping Beauty transposase, we investigate the structural elements that are the most sensitive to mutations, with particular emphasis on protein sectors. We show that sectors are enriched in DNA-binding residues and are highly sensitive to mutations, which cannot be explained by positional conservation. In addition, our analysis suggests that hyperactivity results from the modification of protein-protein interactions, rather than improved protein folding. Wild-type transposases are not optimal for practical use, because they evolved to transpose at relatively low frequency, as high transposition rates harm their host. As a consequence, modifying their activity or insertion patterns through point mutations is of considerable practical importance, and our results may aid their optimization by identifying mutations that are likely to result in transposases with reduced transposition rate.

## RESULTS

**Determination of the tertiary structure of SB transposase and protein core.** The amino acid sequence of the Sleeping Beauty transposase was obtained from Ivics et al.<sup>19</sup>. Experimentally determined protein structures are available for the DDE domain of the transposase<sup>23</sup> and the N-terminal HTH motif of the DNA-binding domain<sup>24</sup>, but not for the entire transposase. Thus we predicted the tertiary structure of Sleeping Beauty with the I-TASSER molecular modeling platform<sup>25,26</sup>, which uses threading and also *ab-initio* modeling for structure prediction. Additionally, we used the coordinates of the existing experimental structures (see above) as constraints (Figure S1A). Due to the availability of high quality templates, a high quality structure prediction was possible: the estimated template modeling (TM) score<sup>27</sup> of the predicted tertiary structure with an experimentally determined structure is 0.86 ( $\pm$  0.1). Models of

this quality can be successfully used in mutagenesis studies and stability analyses<sup>28</sup>. The most similar structure in the Protein Data Bank (PDB, <http://www.rcsb.org>) to the predicted structure (supplementary SB.pdb file) is the Mos1 transposase<sup>29</sup>, which was also the highest ranking template used by I-TASSER (see Figure S1B and C for structural alignments between the Mos1 transposase and the predicted structure, and Figure S2 for a Ramachandran plot of the predicted SB transposase using PROCHECK<sup>30</sup>). Transposases typically function in a dimeric<sup>29,31</sup> or tetrameric enzyme complex<sup>32,33</sup> (and the N-terminal domain of SB was reported to be able to form tetramers *in vitro*<sup>32</sup>), but the high structural and mechanistic similarity of the monomer to Mos1 strongly suggests that the active core unit of the complex is a very similar dimer as the one seen for Mos1. (Nevertheless tetramers may exist and may even be the functional state, for example during assembly.) Thus the monomer produced by I-TASSER was used to build a dimer, using the Mos1 (3HOT) transposase as a template (Figure S1D.) DNA nucleotides were replaced with Chimera<sup>34</sup>, to match the inverted repeats of SB; next the SB transposases were superposed over the Mos1 dimer (3HOT) with TMalign<sup>35</sup>, followed by correction of clashes and minimization. Severe atomic overlaps (e.g. rings penetrated by other groups) in the initial complex model were manually corrected. The model was then subjected to energy minimization *in vacuo* by the steepest descent algorithm in GROMACS5<sup>36</sup> using the CHARMM27 force field. The minimization converged to machine precision with no remaining overlaps between atoms. Visualizations of the protein structures were made with Chimera.

As buried residues in proteins are known to be less tolerant of mutations than exposed residues<sup>37</sup>, we determined relative solvent accessibility of each residue of the structure with DSSP<sup>38</sup> (see methods). The 75 residues with relative solvent accessibility  $\leq 0.1$  were assumed to form the protein core. Residues that take part in protein-protein interactions were determined using the difference in solvent accessible surface areas (SASA) of the monomeric and dimeric form of SB: all residues that have different SASA in the dimer and monomer were assumed to take part in protein-protein interactions. DNA-protein interactions were determined with the SNAP tool of the 3DNA package<sup>39</sup>.

**Identification of sectors of the SB transposase.** To identify sectors in SB, multiple alignments were made with three different state-of-art tools: *muscle*<sup>40</sup>, *probcons*<sup>41</sup> and *mafft*<sup>42</sup> (see methods). Using the three alignments, statistical coupling analyses were performed to identify protein sectors, with the method described by Halabi et al<sup>1</sup>, using a modified MATLAB script provided by the same study. SCA tests whether the conservation of an amino acid at any position in the sequence alignment is correlated with the conservation of any other residue of the protein<sup>4</sup>, i.e. identifies residues that coevolve. First, it builds a weighted correlation matrix of coevolving amino acids for all residues in the alignment (Figure 1A), and this matrix is subsequently cleaned from statistical noise with a randomization method<sup>1</sup>. The analysis of eigenvalue spectra identified three significant eigenvalues for all three alignments (after the exclusion of the largest one), indicating that there might be up to three sectors in the protein. However, after examining residue weights along eigenvectors 2-4 we could identify only two sectors along eigenvector 2 (see Figure S3) that had similar residue compositions irrespectively of the alignment used (Table S3-S4). Due to the different spatial pattern of the residue weights of the three alignments, attempts to identify a third sector resulted in a poorly defined sector, which had different residues depending on the alignment used, and was also strongly correlated with the other two sectors. In consequence, we use only the two sectors that could be consistently defined in all three alignments, which together contain 72-78 residues, depending on the alignment. The cleaned SCA matrices of all three alignments show that the two sectors are essentially independent (Figure 1B), i.e. the correlations between the residues of a sector are much stronger than the correlations between sectors.

The location of the sectors in the transposase structure is somewhat different from the pattern observed in smaller proteins<sup>1</sup> (Figure 1E-F). Residues of both sectors are located in more than one conserved domain, and in the case of the second sector, residues are present in all three Pfam<sup>43</sup> domains of the transposase (Figure 1C,F), indicating that the division to conserved domains does not strictly correspond with the units of the protein that actually coevolve. Sectors (but also conserved residues) are enriched in DNA-binding residues: their fraction is 29%, as opposed to the 17% observed for the entire protein ( $p < 0.05$  for all three alignments, tests of proportions), but there is no significant difference between the two sectors. The residues of sectors are physically less tightly connected than in most small proteins examined so far, which may arise from the low sequence conservation of the

alignments: inaccuracies in the alignments due to the low sequence similarity result in noise, which reduces correlations among residues, and in consequence SCA may fail to detect certain residues as sector residues. To a lesser degree, minor inaccuracies in the transposon sequences themselves may contribute to such noise, as many transposon sequences – including Sleeping Beauty – are reconstructions of extinct repeats.

**The dependence of transposition rate on sectors, protein core and conservation.** To examine the effect of different residues and protein regions on transposition rate, we used transposition rate measurements of 286 SB mutants, which represent a compilation of all Sleeping Beauty point mutations known to us and also unpublished mutants (see Methods). The distribution of 286 point mutations is approximately uniform across the SB transposase sequence (Figure 2A); however their amino acid distribution is not, as the majority of mutants were alanine replacements (Figure S4). In general, the transposition rates of mutants vary significantly, from completely inactivating the transposase to significantly increasing the transposition rate (Figure 2A). The location of the residues in the protein structure have a large influence on their effect: mutations in protein sectors, conserved residues ( $D > 0.5$ , see methods) and the protein core result in a significantly larger reduction in transposition rate in comparison with the residues that do not belong to any of these groups (Figure 2B, both sectors, conserved residues and the core are significantly different from other residues,  $p < 0.05$ , pairwise comparisons with Mann-Whitney U tests).

Sectors represent an extension of the traditional concept of conservation, and there is significant overlap between residues that are part of a sector and also have high positional conservation ( $D > 0.5$ ). Recently it has been questioned whether the effect of sectors on transposition rate is independent from the effect of conservation<sup>2</sup>. To test this, we split sector and conserved residues into three groups: sector residues with low conservation ( $D < 0.5$ ), sector residues with high conservation, and conserved residues that are not part of a sector. The comparison of these groups with residues that are neither part of sectors, nor the protein core, and are also not conserved (“other” residues, Figure 3) indicates, that the effect of sectors on transposition rate is not simply due to positional conservation, as the three groups are significantly different from the “other” residues ( $p < 0.05$  for all comparisons except “conserved only” of the



mafft alignment, Fisher post-hoc tests, ANOVA on log transformed transposition rates). There is no significant difference between non-conserved sector and non-sector conserved residues ( $p > 0.05$  in all three alignments, Fisher post-hoc tests, ANOVA).

**The effect of mutations on protein stability.** Most proteins can function only in a narrow range of folding energies<sup>44</sup>, as unstable proteins may not fold properly and very stable ones may be too rigid to perform their functions. Mutating a residue in a protein can have significant effects on its overall stability ( $\Delta G$ , the free energy of unfolding) and function, thus we tested whether the differences in transposition rate between the sectors, conserved residues and core of the protein and other residues are caused by their effect on protein stability, measured as the difference of the predicted folding energy ( $\Delta\Delta G$ ) between the wild type SB transposase and the mutants. The analysis shows that mutations in sector, conserved and core residues usually have a destabilizing effect on the structure ( $\Delta\Delta G > 0$ ,  $p < 0.05$ , t-tests; Figure 4A).

Although three conserved domains were identified in the SB sequence, an analysis of the flexibility of the structure with the PiSQRD tool<sup>45</sup> and also recent analyses of the Mos1 and SB transposase<sup>24,46</sup> indicate that the structure can be split into two large regions; the relatively flexible N-terminal part of the protein containing the DNA binding HTH-domains (residues 1-120), and a rigid, globular region (residues 121-340) containing the DDE domain (Figure 4B, Figure 1C). Mutations have different effects on folding energies in these two regions; while we detected a clear negative correlation ( $p < 0.001$ ,  $R = -0.51$ ) between transposition rate and  $\Delta\Delta G$  (Figure 4C) in the globular part of the protein, there is no such relationship ( $p = 0.95$ ,  $R = 0.0049$ ) in the N-terminal region containing the HTH domains (Figure 4D).

Next we tested whether mutants in the two regions have different effects on the transposition rate of SB, and we found that the two regions are markedly different. In the flexible part of the protein mutants of sector, conserved and core residues do not differ significantly from the remaining residues ( $p > 0.05$  for all comparisons, Mann-Whitney U tests, Figure 5A), while in the region containing the DDE domain there is a highly significant difference ( $p < 0.001$  for all comparisons, Mann-Whitney U tests, Figure 5B). Additionally, 50% of the mutants of “other” residues have higher transposition rates than the wild type.

As the location of mutations has a significant effect on the free energy of folding, and in the DDE domain  $\Delta\Delta G$  is correlated with transposition rates (Figure 4D), we tested whether the effect of sectors remains significant if we remove the effect of  $\Delta\Delta G$  on transposition rate, i.e. we adjust all rates to  $\Delta\Delta G = 0$ . The results show that the corrected transposition rates are still highly significantly different from other residues ( $p < 0.05$  for all comparisons, Mann-Whitney U tests, Figure 5C), thus the biological effect of sectors and conserved residues cannot be explained with their effect on  $\Delta\Delta G$  alone.

**The effect of protein-protein and protein-DNA interactions on transposition rate.** Transposases typically form protein complexes during transposition, and recent studies on mariner transposases related to Sleeping Beauty (Hsmar1 and Mos1) indicate that mutants that disrupt allosteric communication within its dimer are characterized by increased activity<sup>47–49</sup>. In particular, almost all mutants of the conserved WVPHEL motif (except P and E) of Hsmar1 transposon were hyperactive<sup>48</sup>, most likely due to lowering the kinetic barrier to synapsis<sup>50,51</sup>. Our findings suggest that the mechanism that causes hyperactivity in SB may be comparable to Hsmar1, and probably involves the modification (or disruption) of protein–protein interactions (although the WVPHEL motif is not conserved in the SB transposase). This hypothesis is also consistent with the observation that the relationship between transposase concentration and SB activity is similar to Hsmar1<sup>51</sup>. We tested whether mutants of residues taking part in protein-protein and DNA-protein interactions have different transposition rates than other residues at the protein surface. In general, when outliers are excluded, mutants of both protein and DNA-interacting residues have significantly lower transposition rates than other residues at the surface (Figure 6,  $p < 0.05$ , Kruskal-Wallis test). However, all but two of the hyperactive mutants (with 300% or higher activity) are located at the protein surface, and none are present in the core of the DDE domain. Of the 12 hyperactive surface mutations, four are in the protein-protein interfaces of the dimer (including the most active mutant), and none are in DNA-protein interfaces (see Figure 7 and supplementary Chimera visualization). Since the SB transposase can probably also form tetramers during transposition<sup>32</sup>, there are probably more residues that take part in protein-protein interactions, suggesting that modification of interactions might be a key factor responsible for hyperactivity.

## DISCUSSION

We performed an analysis of protein sectors in a relatively large, multidomain protein with a complex tertiary and quaternary structure, and attempt to predict the effect of mutations on transposition rate, based on their location and effect on protein stability. Although sector identification depends on the alignment used, we could identify two sectors in the SB transposase, regardless of the alignment method. Most previous studies focused on smaller, single-domain proteins<sup>1,7,52</sup>, and one study<sup>8</sup> identified a sector that spans two domains; our analysis indicates that sectors can span multiple conserved domains of a protein (Figure 1E-F), and, in the case of SB, are enriched in DNA binding residues. **There may be at least two explanations for the observation that sectors residues are present in more than one domain: first, in some stages of transposition these residues may be in physical contact. Second, since sector identification is a purely statistical procedure which searches for coevolving residues in the entire protein sequence, in the case of two (or multi) domain proteins where both domains are necessary for the protein to function, coevolution between the domains is highly likely, and sectors that are confined to a single domain are probably present only in domains that are essentially independent.**

A significant effect of mutations on transposition rate could be demonstrated in sectors, the protein core, conserved residues, protein-protein and protein-DNA interface: mutating these residues typically resulted in transposases with low transposition rates. Recently Teşileanu et al. suggested that depending on the method used for sector identification, any biological effect of the first sector may be the consequence of sequence conservation alone<sup>2</sup>. Since we used the method of Halabi et al.<sup>1</sup> for sector identification, which does not use the first eigenvector of the SCA correlation matrix, their concerns do not apply for our results. However, as a significant fraction of sector residues are conserved, we also analyzed sector and conserved residues independently, which shows that mutations of not conserved sector residues have a similar effect on transposition rate as mutations of conserved but non-sector residues (Figure 3), thus the biological functions of sectors cannot be explained with conservation alone.

In comparison with smaller proteins<sup>5</sup>, the influence of sectors on protein function appears to be more complicated, and depend on the tertiary structure. In the globular part of the protein, we could detect a clear effect of sectors, conservation and core on transposition rate, even when the effect of the free energy of folding was excluded (Figures 5). However, in the flexible part of the protein containing the

HTH-domains we found no effect of sectors, nor a correlation between transposition rate and  $\Delta\Delta G$  (Figure 4), which indicates that further studies are needed to evaluate the importance of sectors in non-globular (including disordered and coiled coil) proteins.

While we didn't find a "recipe" for making hyperactive mutants of SB, our analysis allows prioritizing residues for targeted mutagenesis. Half of the residues in the DDE domain that are not part of sectors, conserved residues or protein core have increased transpositional activity. In addition, 12 out of the 14 hyperactive mutants (mutants with at least 3x increased activity compared to the wild type) are located in the protein surface, and 4 of them are in the protein-protein interfaces of the dimer, suggesting that similarly to the Hsma1, the disruption of self-regulating protein-protein interactions may be an important factor in generating hyperactive mutants. In contrast, no hyperactive mutants are present in DNA-protein interfaces or in the buried residues (core) of the DDE domain. Since mutations of these regions typically strongly reduce the rate of transposition, this suggests that despite the fact that SB is a reconstructed sequence and it is most likely inaccurate to some degree, both DNA binding and folding are close to optimal.

## MATERIALS AND METHODS

**Identification of SB homologs and making of multiple alignments.** Transposase sequences homologous to Sleeping Beauty were identified in the 6-frame translated RepBase database (v17.12)<sup>53</sup>, the main database of eukaryotic transposable elements, using the jackhmmer tool of the HMMER 3.0 package<sup>54</sup>, with bit score cutoff 27. We excluded from the hits all matches that show homology only to a short fragment of SB, and kept only those hits that span at least from residues 50 to 290 of the SB transposase, thus covering more than 70% of the sequence. Next, to remove sequences with high similarity (>90%), the homologous sequences were clustered with uclust<sup>55</sup>. The determination of protein sectors depends on multiple sequence alignments, but in the case of SB the average pairwise sequence similarity between the homologous sequences is low (19%), and in this low range of sequence similarity only approximately 50-80% of the residues can be aligned correctly with current methods<sup>56</sup>. This means that the choice of the aligner may influence the results significantly (i.e. the determination of sector and conserved residues), and to estimate the biases introduced by different alignment methods, we used

three different alignment tools: *muscle*<sup>40</sup>, *probcons*<sup>41</sup> and *mafft*<sup>42</sup>. After aligning the sequences, the alignments were trimmed to the 340 residues of SB transposase, i.e. we removed all columns with gaps in the SB sequence. All three alignments are available for download as Supplementary Data.

**Determination of conservation and SCA calculations.** Conservation ( $D$ , Kullback-Leibler entropy) at any given position of the sequence was defined as the divergence of the observed frequency from the background frequency of the most frequent residue at the position, and was calculated with the following equation:  $D = f \ln(f/q) + (1-f) \ln[(1-f)/(1-q)]$ <sup>1</sup>, where  $f$  denotes the frequency of the amino acid at a given position of the sequence, and  $q$  represents its background frequency. We used the same background frequencies as in<sup>1</sup>, and conserved residues were defined as residues with  $D > 0.5$ . Both for SCA and  $D$  calculations, we excluded all positions from the alignments, where the frequency of gaps was higher than 30%. SCA calculations (calculating the correlation matrix, spectral cleaning, randomization of the alignments) were performed with a modified Matlab script provided by the Halabi et al.<sup>1</sup>. Sectors were determined by a visual examination of residue weights of eigenvectors 2-4 (see Halabi et al. for details), sector 1 was defined as residues with weights  $< -0.05$  along eigenvector 2, sector 2 as residues with weights  $> 0.05$  along eigenvector 2 (see Figure S3).

**Construction of SB mutants and determining their transposition rate.** The mutants were partly obtained from published studies<sup>20,57-60</sup>, and partly (~80 mutants) represent unpublished material. Site-directed mutagenesis of the transposase gene was done by PCR following the QuikChange (Stratagene) principle of site-directed mutagenesis. The mutants were tested against the corresponding wild-type SB transposase in cell-based transposition assays, as originally described by Ivics et al.<sup>19</sup>.

**Stability calculations and in-silico mutagenesis.** The free energy of unfolding ( $\Delta G$ ) of the SB transposase, and its changes were calculated with the FoldX tool (version 4)<sup>61,62</sup>, using the predicted structure of the SB complex. First, the structure produced by I-TASSER was optimized with the RepairPDB function to correct torsion angles, van der Waals clashes and total energies. Next we calculated the effect of the mutations on the  $\Delta G$  of the structure for the 286 mutants. The difference in  $\Delta G$

between the 'wild type' SB and its mutants is given as  $\Delta\Delta G$ ; its positive values indicate destabilizing, negative values stabilizing mutations.

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## SUPPLEMENTARY MATERIAL

Supplementary material is available in the online version of the paper: Supplementary figures 1-4; Supplementary Tables 1-4; the predicted structure of the Sleeping Beauty transposase ([SB.pdb](#)); Chimera visualizations of the transposase dimer showing the hyperactive (300+% activity) mutants ([hyperactive.py](#)) and the mutants marked as "other" ([other.py](#)); and the alignments used to determine protein sectors. A list with 229 SB mutants is available in the attached [mutations.txt](#) file. The remaining 57 mutants are available from Wang Y., Nagy E.E, Pryputniewicz-Dobrinska D. et al.: Regulated Complex Assembly Safeguards the Fidelity of *Sleeping Beauty* Transposition (Submitted).

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## FIGURE LEGENDS

### Figure 1. Identification of sectors and conserved domains in the Sleeping Beauty (SB)

**transposase.** A) Statistical Coupling Analysis (SCA) matrix for the muscle alignment of 289 homologous sequences present in RepBase (+SB). The matrix represents correlations between amino acid frequencies at each position of the alignment, i.e. residue pairs that coevolve. B) Cleaned SCA matrices for three alignments made with *muscle*, *probcons*, and *mafft* aligners, containing the residues of the two sectors. Residues within sectors show correlated evolution, while there is almost no correlation between sectors. C) The transposase contains three Pfam conserved domains; two HTH domains with DNA binding functions, and a DDE domain with endonuclease activity. D) The distribution of conservation scores (*D*) across the sequence. E-F) The location of the two sectors identified with the *muscle* alignment in the tertiary structure of the SB transposase. The sectors are located across secondary structure elements, and are less compact than the ones reported so far, possibly due to the low sequence similarity in the alignments. Both sectors have residues in multiple conserved domains; most notably sector 2, which has residues in all three Pfam domains of the protein. G) The location of the conserved residues (*D* > 0.5, muscle alignment) of the transposase. H) The residues of the protein core. All residues with relative solvent accessibility below 0.1 are highlighted with red.

**Figure 2. Effect of residue location on the transposition rate of SB mutants.** A) The location of the mutations along the SB transposase sequence, and their effect on transposition rate. The 286 mutants are distributed approximately evenly across the sequence; the majority of mutants reduces transposition rate ( $< 100\%$  of SB). None of the Pfam conserved domains show a clear difference from the rest of the sequence. B) The effect of sectors, conserved residues and protein core on transposition rate (median, box: 25%-75%, whiskers: 10%-90%). Mutants in both sectors, conserved residues and residues of the protein core have significantly lower transposition rates than other residues, irrespectively of the aligner used ( $p < 0.05$ , Mann-Whitney U tests).

**Figure 3. The effect of sectors on transposition rate is not a by-product of positional conservation.** Sector and conserved residues were split into three groups: sector residues with low positional conservation ( $D < 0.5$ ), sector residues with high positional conservation, and conserved residues that are not part of any sector. Transposition rates of mutants (median, box: 25%-75%, whiskers: 10%-90%) in all three groups are significantly different from mutants in other residues ( $p < 0.05$  for all comparisons, Fisher post-hoc tests, ANOVA on log transformed transposition rates), and there is no significant difference between the mutants of non-conserved sector and conserved but non-sector residues ( $p > 0.05$  in all three alignments, Fisher post-hoc tests).

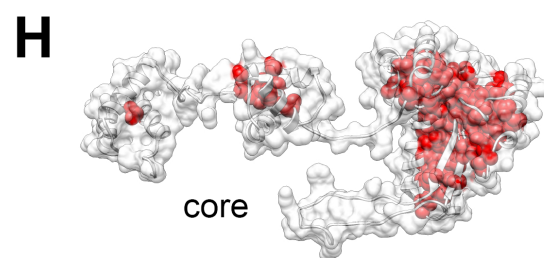
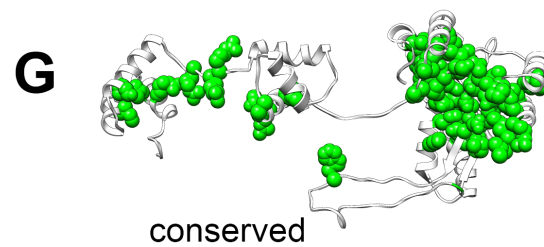
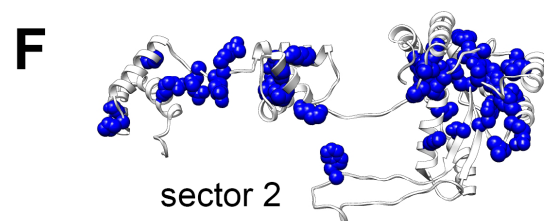
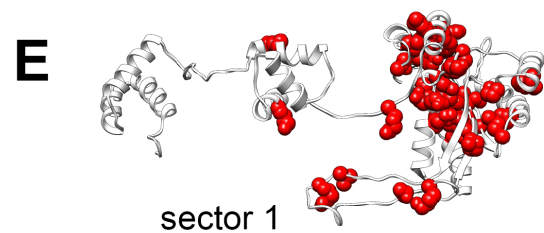
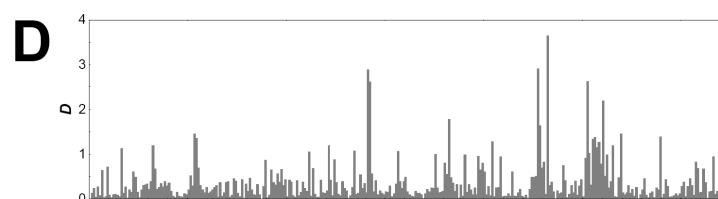
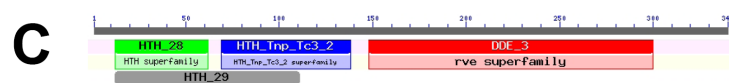
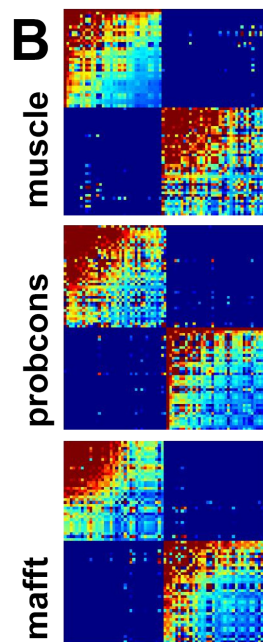
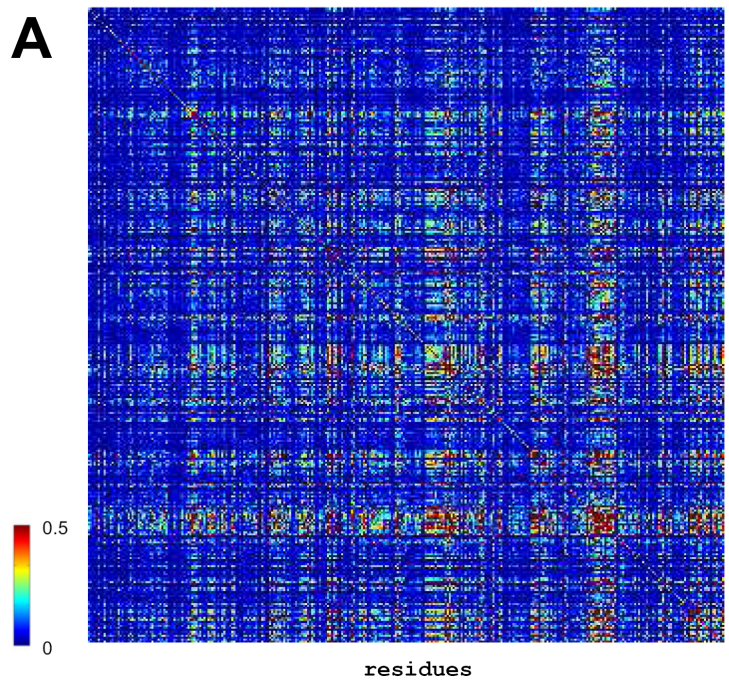
**Figure 4. The effect of mutations on the change of the free energy of unfolding** (median, box: 25%-75%, whiskers: 10%-90%). A) Mutations in sectors and the core are significantly more destabilizing ( $\Delta\Delta G > 0$ ) than mutants of other residues ( $p < 0.05$  for all comparisons, t-tests). B) **The monomer of SB.** The flexible N-terminal arm of the protein containing the HTH domains (residues 1-120) is indicated with white, the globular part (residues 121-340), which contains the DDE domain, with gray. C) In the flexible arm the effect of mutations on  $\Delta\Delta G$  is not correlated with transposition rate ( $p = 0.95$ ). D) In the globular region we find a significant negative correlation ( $p < 0.001$ ,  $R = -0.51$ ) between  $\Delta\Delta G$  and transposition rate.

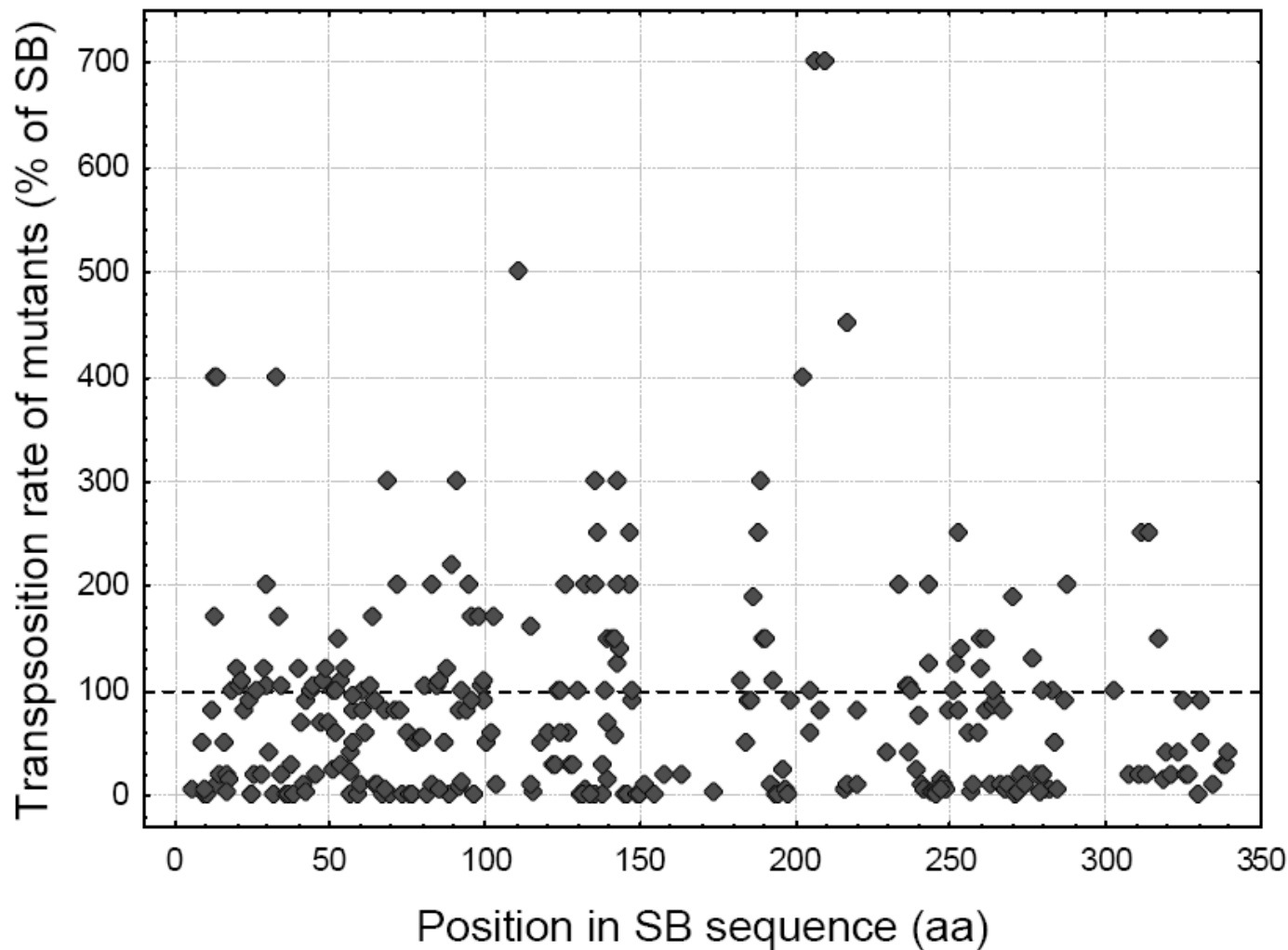
**Figure 5. The effect of residue location on transposition rate, in the two regions of the protein** (median, box: 25%-75%, whiskers: 10%-90%). A) In the HTH-region, mutants of sectors, conserved or

buried residues are not significantly different from the remaining mutants ( $p > 0.05$  for all comparisons, Mann-Whitney U tests). B) In the DDE domain the differences are highly significant ( $p < 0.05$  for all comparisons, Mann-Whitney U tests), even after correcting for the different effects of free energy of folding (C). Note that in the DDE domain 50% of mutants of “other” residues are characterized with higher activity than the wild type SB.

**Figure 6. The effect of DNA-protein and protein-protein interactions on transposition rate.** When outliers are excluded, mutants of residues interacting with DNA (“DNA”) and the other SB chain (“PPI”) have significantly lower transposition rates ( $p < 0.05$ , Mann-Whitney U test) than other residues located at the surface ( $RSA > 0.1$ ). Surprisingly, 12 of the 14 hyperactive mutants (outliers, 300+% activity) are also located in the protein surface, and none in the DNA binding regions, suggesting that the modification of protein-protein interactions might be responsible for their dramatically increased activity. (Outliers with identical transposition rates were shifted by 10%, for visibility)

**Figure 7. The location of the 14 hyperactive mutations in the SB dimer.** Yellow residues represent mutants in protein-protein interfaces, red residues other mutants. As SB probably forms also a tetramer in certain phases of transposition, the number of residues taking part in PPIs is probably higher. (See also the supplementary “hyperactive.py” Chimera file.)



**A****B**